OTIC FILE COPY

AD-A228 684

Quantitative Assessment and Reduction of Long-term Autoradiographic Background

RICHARD K. TRAUB, LUCINDA FAMOUS, RAJESH KRISHNAN, and KAREN R. OLSON

Neurotoxicology Branch, Pathophysiology Division, United States Army Medical Research Institute of Chemical Defense, Aberdeen Proving Ground, Maryland 21010-5425.

Received for publication June 20, 1989 and in revised form November 22, 1989; accepted December 1, 1989 (9A1724).

Quantitative autoradiography can measure distribution patterns in an animal exposed to radiolabeled compounds. A comparison of autoradiographs of rat brain containing low levels of ¹⁴C showed that a highly variable background signal had been produced. This resulted in several overexposed autoradiographs which could not be quantitatively compared. The background, believed to be produced by light emanating from the phosphor coating in the X-ray cassette, was a major impediment because it hindered correct analysis of the specimen. This article details our experiments demonstrative.

strating the sources of variance contributing to background and offers methods for its reduction. We found that placement of black polyethylene plastic between the slides and phosphor in the X-ray film cassette minimized autoradiographic background and effectively eliminated the effects caused by inherently different levels of radioactivity in the glass slides. (J Histochem Cytochem 38:581-583, 1990) KEY WORDS: Quantitative: Autoradiography, Phosphor; Background; Variance reduction: Rat; Central nervous system.

Introduction

Quantitative autoradiography provides solutions to several technical problems encountered in the field of biology (1-4). Actions of endogenous and exogenous compounds can be followed through the course of distribution, binding, metabolism, and elimination (2, 9). In addition, the kinetics of these processes and the associated functional implications of compound interactions can be determined (5-7). Track-tracing autoradiography can define submicrometer binding loci (8).

In our work, we used ^{14}C -labeled soman, a powerful organophosphorous anti-cholinesterase (specific activity 56 mCi/mmol) at a subconvulsive dose of 17.3 µg/kg, IM. The objective was to quantitate the kinetics of distribution in the CNS of rats (9). As a result of both the low specific activity of the labeled soman and the extremely low dose, long exposure times were required to produce a readable autoradiographic exposure. We expected soman to accumulate at cholinesterase sites.

We employed image analysis and enhancement techniques to improve signal-to-noise ratios and quantitate results. One problem we encountered was a large variance in background optical density. Our hypothesis was that glass radionuclides were interacting with the autoradiographic film and/or the cassette enhancement phosphor. This report attempts to define the source and the amount of variance and recommends methods for its reduction.

Materials and Methods

Autoradiography. Autoradiographic data were collected from frozen rat brain tissue mounted on glass slides as described elsewhere (9). Three groups of 26 slides each were placed in X-ray film cassettes (Spectroline 8×10 -inch cassette; Spectronics Corp., Westbury, NY) with the tissue apposed by DuPont Lo Dose mammography film (emulsion side facing tissue) and were exposed for 703 days. These same slides were subsequently re-exposed for 736 days under identical conditions and configurations, with the exception that a black polyethylene sheet 0.005 inches thick was placed between the slide and the phosphor coating on the bottom surface of the cassette. An unexposed control film was also developed to measure film contribution to optical density, generating a total of seven films. The films, including the control, were developed according to standardized methods (7).

Image Acquisition and Analysis. A Hamamatsu Image Acquisition System C1440 (Hamamatsu Systems; Waltham, MA) was used. Images were acquired under non-varying light and camera conditions. High acquisition system sensitivity required elimination of all extraneous light.

All images of slides from both of the 2-year studies, controls, and light source image data were analyzed using a custom program (Image Analysis System, v9.9; DAKKRO Corp. Denver CO). The images were acquired as transmitted light data. Reduction of variance in slide images required the application of corrections derived from analysis of light source and clear film data. The slide image data were divided by control film data for all six samples. The divided images were converted to optical density.

After generating the optical densities for all images, the data were evaluated using a program function of the image analysis system that calculated optical density statistics for user-selected portions of the image. Data were collected horizontally across the non-tissue (glass) portion of each image.

DISTRIBUTION STATEMENT A

Approved for public releases

Distribution Unlimited

¹ To whom correspondence should be addressed.

System response linearity to transmitted light was determined using an optical density wedge.

Two-tailed t-tests were used to analyze the data obtained from the images.

Results

The results in Table 1 [images (A) through (F)] clearly show that blocking the light from the phosphot with the plastic essentially eliminated the increased exposure due to phosphot ($p \ge 0.9999$, t-test). In images (A) and (B), it normalized the differences in the two types of slides ($p \ge 0.9999$, t-test). In addition, there are essential differences in the level of signal generated by the two slide producers (A and C).

Regression analysis of optical wedge data provided the equation OD = 0.0129 + 0.1217(X) with a correlation coefficient of 0.999985.

Statistical analysis of the clear control film yielded the following results: mean OD of 0.4731, standard deviation of 0.0029.

An analysis of signal-to-noise ratios for the various combinations of tissues and slide backgrounds showed ratios varying from 6:1 to 18:1 in the data generated from the cassettes with opaque light block to 0.28:1 to 0.62:1 for cassettes with no opaque blocking.

Discussion

The variation observed from both the measuring system and the toradiographic configuration showed that small differences in au ical density could be reliably measured. The differences in opterials used to obtain autoradiographic data, such as the glass males and type of holding cassette, appeared to be the major source of the observed variation detailed below.

System variance (from camera, electronics, and lens nonlinearities) was present in all of the original images. Images from the light source (LS) and light source plus film (LSF) were used to define this variance and the methods for its reduction. In LS and LSF, there was a non-random fluctuation of transmitted light data values resulting from spherical lens astigmatism, which was correctable. There were other minor non-random signal sources from camera electronics. The percentage variation was 33% for both LS and LSF. Therefore, LSF was used to remove variance due to the system from the subsequently measured slide images by division. This process had no effect on the OD values from non-system errors. Using LSF to normalize the slide images reduced system variance from 33% to 2.7%.

The method described above was able to reduce the amount of system variation in the slide images. However, variation between individual glass slides was still evident. The hypothesis was that radioisotopes from the glass slides interacted with the covering of phosphor used to enhance X-rays in the cassette, thus causing light to be produced. The film in the first 2-year study was exposed to this excess light and produced a higher optical density than the subsequent exposures. To block light from the phosphor, a piece of opaque black polyethylene plastic was placed between the glass slides and phosphor.

Analysis of the results from quantitative optical density analysis of six slide/cassette configurations showed three significant trends. First, Table 1 summarizes the results of the OD measurements and shows that there are major differences between cassettes with and

Table 1. Source and quantitative image data

Image ⁴	Slide	N'	Mean optical density (OD)	Standard deviation
			<u></u>	of OD
(A)	C۱	34	0.6202	0.0020
	C_2	39	0.6177	0.0031
	\mathbf{A}_{I}	37	0.5769	0.0024
	\mathbf{A}_2	35	0.5749	0.0020
(B)	C_1	36	0.4984	0.0039
	C_2	33	0.4980	0.0023
	A_1	39	0.4923	0.0018
	\mathbf{A}_2	34	0.4953	0.0022
(C)	A 3	41	0.5195	0.0016
	A_4	42	0.5153	0.0020
	A 5	40	0.5176	0.0020
	A_6	36	0.5194	0.0020
(D)	A,	37	0.5794	0.0027
	A.4	37	0.5760	0.0027
	A5	38	0.5753	0.0036
	A_6	39	0.5739	0.0028
(E)	Α-	38	0.5721	0.0048
	C_3	38	0.6493	0.0052
	A_8	38	0.5729	0.0014
	C4	38	0.6329	0.0022
(F)	A ₂	41	0.4964	0.0019
	C_3	43	0.5122	0.0018
	A_8	39	0.4965	0.0018
	C ₄	40	0.5148	0.0015

Images: (A) Cassette E1 developed without black plastic; (B) cassette E1 developed with black plastic; (C) cassette E2 developed with black plastic; (D) cassette E2 developed without black plastic; (E) cassettes E1-B3 developed without black plastic; (F) cassettes E1-B3 developed with black plastic.

^b A slides from American Scientific; C slides from Corning Glass

'N, number of image pixels analyzed

without black plastic between the cassette phosphor and slides. Second, the results show that there are major differences between the levels of background radiation emitted by the slides tested from two different manufacturers. It should be noted that this observation is only valid for these lot numbers of slides and cannot be generalized to all slides produced by these manufacturers. This was not quantitatively analyzed as a part of this study. However, it was confirmed by visually comparing slides from these two companies in previous studies and observing that not all slides were consistently light or dark across producers. Rather, it seemed to vary as a function of the lot numbers and appeared consistent within a lot rather than across lots. Third, the inclusion of the black plastic eliminates the difference between slides in most instances.

An analysis of signal-to-noise ratios indicated that fractional ratios were produced by the cassette configuration that contained no light block. Fractional signal-to-noise ratios mean that there is more noise than signal, which in this case is physically observed as tissue absorbing light emitted by the phosphor which has been activated by the glass radionuclides. The tissue is actually less optically dense than the background. This situation produces autoradiographic data that are not related in any simple way to radionuclides in tissues. It is a complex function of differential absorbance of transmitted light and tissue radionuclide emission.

1

For long-term autoradiographic exposures, the effect of unwanted background radiation can be reduced by using cassettes without phosphor enhancers and/or phosphor light output blocked by opaque sheets, together with slides of low inherent radioactivity.

Acknowledgment

We wish to thank Mr H. Shafer for technical assistance.

Literature Cited

- 1. Benfenati F, Cimino M, Agnati L, Fuxe K: Quantitative autoradiography of central neurotransmitter receptors: methodological and statistical aspects with special reference to computer-assisted image analysis. Acta Physiol Scand 128:129, 1986
- Biegon A, Rainbow T: Distribution of imipramine binding sites in the rat brain studied by quantitative autoradiography. Neurosci Lett 37:209, 1983
- Carney H, Fiahnenstiel G: Quantification of track and grain density autoradiography and evaluation of 14C loss on preservation. J Plankton Res 9:41, 1987

- 4. Filitney F: Autoradiography. In Bancroft J, Stevens A, eds. Theory and practice of histological techniques. New York, Churchill-Livingstone, 1977, 371
- Katz D, Kimelberg H: Kinetics and autoradiography of high affinity uptake of serotonin by primary astrocyte cultures. J Neurosci 5:1901, 1985
- Keller F, Waser P: Brain pharmacokinetics of centrally acting drugs, a quantitative autoradiographic study. Arch Int Pharmacodyn Ther 267:200, 1984
- 7. Lev H, Nielsen A, Ursin C: Cell kinetics in the epidermis of the toad Bufo bufo (L) after partial hypophysectomy followed by corticotrophin treatment (an autoradiography study). Comp Biochem Physiol 75A: 51, 1983
- 8. Rogers A: Techniques of autoradiography. London, Elsevier, 1973
- 9. Traub K: In vivo distribution of 14C radiolabeled soman ((3,3-dimethyl-2-butoxy)-methylphosphoryl fluoride) in the central nervous system of the rat. Neurosci Lett 60:219, 1985



Access	ion For						
NTIS	GRA&I	V					
DTIC 7							
Unannounced 🔲							
Just11	rication						
Avai	ibution/ lebility	Codes					
:	Avail a	nd/or					
Dist	al						
PV	20						

REPORT D	Form Approved OMB No. 0704-0188								
1a. REPORT SECURITY CLASSIFICATION UNCLASSIFIED	16 RESTRICTIVE MARKINGS								
2a. SECURITY CLASSIFICATION AUTHORITY	3 DISTRIBUTION / AVAILABILITY OF REPORT								
2b. DECLASSIFICATION / DOWNGRADING SCHEDU	Approved for public release;								
4. PERFORMING ORGANIZATION REPORT NUMBE	distribution unlimited 5. MONITORING ORGANIZATION REPORT NUMBER(S)								
P89-019		P89-019							
6a. NAME OF PERFORMING ORGANIZATION	6b. OFFICE SYMBOL (If applicable)	7a. NAME OF MONITORING ORGANIZATION							
US Army Medical Research	(ii applicable)	US Army Medical Research							
Institute of Chemical Defense	SGRD-UV-YN	Institute of Chemical Defense							
6c. ADDRESS (City, State, and ZIP Code)	7b. ADDRESS (City, State, and ZIP Code)								
Aberdeen Proving Ground, MD 21	010_5/25	Aberdeen Proving Ground, MD 21010-5425							
8a. NAME OF FUNDING/SPONSORING	8b. OFFICE SYMBOL								
ORGANIZATION	(If applicable)	9. PROCUREMENT INSTRUMENT IDENTIFICATION NUMBER							
8c. ADDRESS (City, State, and ZIP Code)		10. SOURCE OF FUNDING NUMBERS							
,, ,		PROGRAM	PROJECT	TASK	WORK UNIT				
		ELEMENT NO.	NO.	NO.	ACCESSION NO.				
		3M61102B	61102A	S11	AA				
11. TITLE (Include Security Classification)	duation of Tona	+ a.um							
Quantitative Assessment and Re Autoradiographic Background	auction of roud-	-cerm							
12. PERSONAL AUTHOR(S)									
Traub, Richard K., Famous, Lu	cinda, Krishnan	. Rajesh. and	d Olson. Kar	en R					
13a. TYPE OF REPORT 13b. TIME CO	OVERED	14. DATE OF REPORT (Year, Month, Day) 15. PAGE COUNT							
Open Literature Publ. FROM	то				3				
16. SUPPLEMENTARY NOTATION									
The Journal of Histochemistry 17. COSATI CODES	and Cytochemist: 18. SUBJECT TERMS (
FIELD GROUP SUB-GROUP	Id. SOBJECT TERMS (Continue on levels	e ir riccessory drie						
,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	Quantitative autoradiography								
	guanozou ozvo								
19. ABSTRACT (Continue on reverse if necessary and identify by block number)									
Quantitative autoradiography can me terns in an animal exposed to radiol comparison of autoradiographs of ra levels of ¹⁴ C showed that a highly van nal had been produced. This resulted autoradiographs which could not be query to be produced to be produced ing from the phosphor coating in the major impediment because it hinde the specimen. This article details out 20. DISTRIBUTION/AVAILABILITY OF ABSTRACT	strating the sources of variance contributing to background and offers methods for its reduction. We found that placement of black polyethylene plastic between the slides and phosphor in the X-ray film cassette minimized autoradiographic background and effectively eliminated the effects caused by inherently different levels of radioactivity in the glass slides. (J Histochem Cytochem 38:581-583, 1990) KEY WORDS: Quantitative; Autoradiography; Phosphor; Background; Variance reduction; Rat; Central nervous system.								
UNCLASSIFIED/UNLIMITED SAME AS F	RPT. DTIC USERS	ADJINACI SE	ZI. ABSTRACT SECURITY CLASSIFICATION						
22a. NAME OF RESPONSIBLE INDIVIDUAL	226 TELEPHONE (Include Area Code) 22c. Of	FFICE SYMBOL					
Wade. John W. LTC	301-671-3	2553	SGR	D-IIV-YN					

DD Form 1473, JUN 86

Previous editions are obsolete

SECURITY CLASSIFICATION OF THIS PAGE